=> d his nofil

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(FILE 'HOME' ENTERED AT 11:11:41 ON 05 DEC 2006)
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FILE 'REGISTRY' ENTERED AT 11:11:54 ON 05 DEC 2006
           2199 SEA ABB=ON PLU=ON "PROPARGYL"
L1
L2
                STR
L3
              0 SEA SSS SAM L2
             26 SEA SSS FUL L2
L4
     FILE 'HCAPLUS' ENTERED AT 11:23:40 ON 05 DEC 2006
L5
              2 SEA ABB=ON PLU=ON L4
     FILE 'BEILSTEIN' ENTERED AT 11:23:50 ON 05 DEC 2006
L6
              0 SEA SSS SAM L2
              O SEA SSS FUL L2
L7
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              2 SEA SSS SAM L2
L8
             10 SEA SSS FUL L2
L9
              8 SEA ABB=ON PLU=ON L9/COM
L10
     FILE 'WPIX' ENTERED AT 11:28:12 ON 05 DEC 2006
              0 SEA SSS SAM L2
L11
              1 SEA SSS FUL L2
L12
                D L12
              1 SEA ABB=ON PLU=ON L12/DCR
L13
                SEL L12 SDCN
L14
              1 SEA ABB=ON PLU=ON RAC5T3/DCN
                SEL L12 DCSE
              0 SEA ABB=ON PLU=ON 800257-0-0-0/DCRE
L15
              1 SEA ABB=ON PLU=ON L13 OR L14
                D COST
                D HITSTR
                D COST
     FILE 'HCAPLUS, MEDLINE, EMBASE, BIOSIS' ENTERED AT 11:42:45 ON 05 DEC 2006
            228 SEA ABB=ON PLU=ON GRAMMENOS W?/AU
L17
L18
            380 SEA ABB=ON PLU=ON GROTE T?/AU
            77 SEA ABB=ON PLU=ON BLETTNER C?/AU
L19
            148 SEA ABB=ON PLU=ON GEWEHR M?/AU
L20
            120 SEA ABB=ON PLU=ON GYPSER A?/AU
L21
         5036 SEA ABB=ON PLU=ON MULLER B?/AU
253 SEA ABB=ON PLU=ON RHEINHEIMER J?/AU
L22
L23
            691 SEA ABB=ON PLU=ON SCHAFER P?/AU
L24
            11 SEA ABB=ON PLU=ON SCHWOGLER A?/AU
L25
L*** DEL
            0 S TRNO J?/AU
L*** DEL
            176 S L17 AND L18
L*** DEL
            1 S L26 AND L25
                D BIB
            275 SEA ABB=ON PLU=ON TORMO J?/AU
L26
·L27
           99 SEA ABB=ON PLU=ON GOTZ N?/AU
L28
           1238 SEA ABB=ON PLU=ON LORENZ G?/AU
            842 SEA ABB=ON PLU=ON AMMERMANN E?/AU
            479 SEA ABB=ON PLU=ON STRATHMANN S?/AU
L30
L31
            235 SEA ABB=ON PLU=ON STIERL R?/AU
            790 SEA ABB=ON PLU=ON (L17 AND (L18 OR L19 OR L20 OR L21 OR L22
L32
                OR L23 OR L24 OR L25 OR L26 OR L27 OR L28 OR L29 OR L30 OR
                L31)) OR (L18 AND (L19 OR L20 OR L21 OR L22 OR L23 OR L24 OR
                L25 OR L26 OR L27 OR L28 OR L29 OR L30 OR L31)) OR (L19 AND
```

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(L20 OR L21 OR L22 OR L23 OR L24 OR L25 OR L26 OR L27 OR L28
                OR L29 OR L30 OR L31)) OR (L20 AND (L21 OR L22 OR L23 OR L24
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                (L30 AND L31)
L33
             2 SEA ABB=ON PLU=ON L32 AND PHENETHY? AND ?ACRYLAMID?
              2 SEA ABB=ON PLU=ON (L17 OR L18 OR L19 OR L20 OR L21 OR L22 OR
L34
                L23 OR L24 OR L25 OR L26 OR L27 OR L28 OR L29 OR L30 OR L31)
                AND PHENETHY? AND ?ACRYLAMID?
            216 SEA ABB=ON PLU=ON L17 AND (L18 OR L19 OR L20 OR L21 OR L22
L35
                OR L23 OR L24 OR L25 OR L26 OR L27 OR L28 OR L29 OR L30 OR
L36
            283 SEA ABB=ON PLU=ON L18 AND (L19 OR L20 OR L21 OR L22 OR L23
                OR L24 OR L25 OR L26 OR L27 OR L28 OR L29 OR L30 OR L31)
L*** DEL
            66 S L19 AND L21-31
             66 SEA ABB=ON PLU=ON L19 AND (L20 OR L21 OR L22 OR L23 OR L24
L37
                OR L25 OR L26 OR L27 OR L28 OR L29 OR L30 OR L31)
L38
            136 SEA ABB=ON PLU=ON L20 AND (L21 OR L22 OR L23 OR L24 OR L25
                OR L26 OR L27 OR L28 OR L29 OR L30 OR L31)
L39
            102 SEA ABB=ON PLU=ON L21 AND (L22 OR L23 OR L24 OR L25 OR L26
                OR L27 OR L28 OR L29 OR L30 OR L31)
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L40
                OR L28 OR L29 OR L30 OR L31)
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L41
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            14 SEA ABB=ON PLU=ON L24 AND (L25 OR L26 OR L27 OR L28 OR L29
L42
                OR L30 OR L31)
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L43
               OR L31)
L44
             0 SEA ABB=ON PLU=ON L26 AND (L27 OR L28 OR L29 OR L30 OR L31)
            28 SEA ABB=ON PLU=ON L27 AND (L28 OR L29 OR L30 OR L31)
L45
L46
           605 SEA ABB=ON PLU=ON L28 AND (L29 OR L30 OR L31)
L47
           357 SEA ABB=ON PLU=ON L29 AND (L30 OR L31)
L48
           195 SEA ABB=ON PLU=ON L30 AND L31
L*** DEL
           318 S (L35 AND L36-48) OR (L36 AND L37-48)
           505 SEA ABB=ON PLU=ON (L35 AND (L36 OR L37 OR L38 OR L39 OR L40
L49
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                (L47 AND L48)
                D SCA L33
L50
           461 SEA ABB=ON PLU=ON L49 AND FUNGICID?
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STRUCTURE SEARCH

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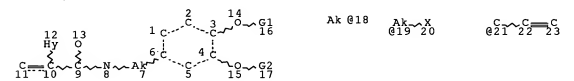
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FILE COVERS 1907 - 5 Dec 2006 VOL 145 ISS 24 FILE LAST UPDATED: 4 Dec 2006 (20061204/ED)

New CAS Information Use Policies, enter HELP USAGETERMS for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

=> d que 15 L2



VAR G1=18/19/21 VAR G2=18/19 · NODE ATTRIBUTES: CONNECT IS E2 RC AT CONNECT IS E2 RC AT CONNECT IS E3 RC AT 9 CONNECT IS E1 RC AT 13 CONNECT IS E1 RC AT DEFAULT MLEVEL IS ATOM 7 12 18 19 MLEVEL IS CLASS AT GGCAT IS LIN LOC SAT AT IS UNS AT GGCAT 12 DEFAULT ECLEVEL IS LIMITED

STR

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 23

STEREO ATTRIBUTES: NONE

L4 26 SEA FILE=REGISTRY SSS FUL L2
L5 2 SEA FILE=HCAPLUS ABB=ON PLU=ON L4

=> fil marpat

FILE 'MARPAT' ENTERED AT 11:59:02 ON 05 DEC 2006
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FILE CONTENT: 1961-PRESENT VOL 145 ISS 22 (20061201/ED)

SOME MARPAT RECORDS ARE DERIVED FROM INPI DATA FOR 1961-1987

MOST RECENT CITATIONS FOR PATENTS FROM MAJOR ISSUING AGENCIES (COVERAGE TO THESE DATES IS NOT COMPLETE):

US 20060234956 19 OCT 2006
DE 102005016345 12 OCT 2006
EP 1710237 11 OCT 2006
JP 2006282618 19 OCT 2006
WO 2006108879 19 OCT 2006
GB 2424583 04 OCT 2006
FR 2884252 13 OCT 2006
RU 2284857 10 OCT 2006
CA 2500558 10 SEP 2006

Expanded G-group definition display now available.

=> d que 110

L2

STR

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2 14 G1 16 Hy 0 6 c 4 c 15 T5 T7
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VAR G1=18/19/21 VAR G2=18/19 NODE ATTRIBUTES: CONNECT IS E2 RC AT CONNECT IS E2 RC AT CONNECT IS E3 RC AT CONNECT IS E1 RC AT 13 CONNECT IS E1 RC AT DEFAULT MLEVEL IS ATOM MLEVEL IS CLASS AT 7 12 18 19 IS LIN LOC SAT AT GGCAT IS UNS AT 12 GGCAT

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RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 23

DEFAULT ECLEVEL IS LIMITED

STEREO ATTRIBUTES: NONE

L9 10 SEA FILE=MARPAT SSS FUL L2

L10 8 SEA FILE=MARPAT ABB=ON PLU=ON L9/COM

=> fil wpix FILE 'WPIX' ENTERED AT 11:59:13 ON 05 DEC 2006 COPYRIGHT (C) 2006 THE THOMSON CORPORATION

FILE LAST UPDATED: 4 DEC 2006 <20061204/UP>
MOST RECENT THOMSON SCIENTIFIC UPDATE: 200678 <200678/DW>
DERWENT WORLD PATENTS INDEX SUBSCRIBER FILE, COVERS 1963 TO DATE

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STR

VAR G1=18/19/21 VAR G2=18/19 NODE ATTRIBUTES: CONNECT IS E2 RC AT CONNECT IS E2 RC AT CONNECT IS E3 RC AT 9 CONNECT IS E1 RC AT 13 CONNECT IS E1 RC AT DEFAULT MLEVEL IS ATOM MLEVEL IS CLASS AT 7 12 18 19 IS LIN LOC SAT AT GGCAT IS UNS AT ~12 GGCAT DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

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STEREO ATTRIBUTES: NONE

L12 1 SEA FILE=WPIX SSS FUL L2

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8 DUP REM L5 L10 L16 (3 DUPLICATES REMOVED)

ANSWERS '1-2' FROM FILE HCAPLUS ANSWERS '3-8' FROM FILE MARPAT

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L51 ANSWER 1 OF 8 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 1

ACCESSION NUMBER:

2003:796663 HCAPLUS Full-text

DOCUMENT NUMBER:

139:292160

TITLE:

Preparation of N-(2-phenylethyl)acrylamides as

agricultural fungicides

INVENTOR(S):

Grammenos, Wassilios; Grote, Thomas; Blettner, Carsten; Gewehr, Markus; Gypser, Andreas; Mueller, Bernd; Rheinheimer, Joachim; Schaefer, Peter; Schwoegler, Anja; Tormo i Blasco, Jordi; Goetz, Norbert; Lorenz, Gisela; Ammermann, Eberhard;

Strathmann, Siegfried; Stierl, Reinhard

PATENT ASSIGNEE(S):

BASF Aktiengesellschaft, Germany

SOURCE:

PCT Int. Appl., 53 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.				KIND		DATE		APPLICATION NO.				DATE					
	WO 2003082822			A1 20031009				WO 2002 ED2010						20020227			
WO							WO 2003-EP3212					20030327					
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		PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	TJ,	TM,	TN,	TR,	TT,	TZ,

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                                20031013
                                            AU 2003-216893
                                                                    20030327
     EP 1492768
                                            EP 2003-712104
                          A1
                                20050105
                                                                    20030327
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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                                            US 2003-509112
     US 2005181948
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                                                                    20030327
PRIORITY APPLN. INFO.:
                                             DE 2002-10214177
                                                                    20020328
                                             WO 2003-EP3212
                                                                 W 20030327
                         MARPAT 139:292160
OTHER SOURCE(S):
GI
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R1 O OR3

Title compds. [I; R1, R2 = H, halo, C1-4 (halo)alkyl, C1-4 (halo)alkoxy, C3-10 cycloalkyl; R3 = C1-4 (halo)alkyl, propargyl, C3-4 alkenyl, CH2C.tplbond.CCRaRbRc; Ra, Rb = H, Me; Rc = H, C1-4 alkyl; R4 = Me, haloalkyl; Het = 5-6 membered (fused) (substituted) heterocyclyl], were prepared Thus, 1.28 g (2E)-N-[2-(3,4-dimethoxyphenyl)ethyl]-4-methyl-2- (tributylstannyl)-2-pentenamide (preparation given) in DMF was stirred with 2-bromo-5-trifluoromethylpyridine, Pd(PPh3)4, and Cu2I2 over night at room temperature to give 0.5 g (2Z)-N-[2-(3,4-dimethoxyphenyl)ethyl]-4-methyl-2-[5-(trifluoromethyl)-2-pyridinyl]-2-pentenamide. Several I at 250 ppm gave 95-100% control of Botrytis cinerea on pepper leaves.

IT 609341-62-0P 609341-63-1P 609341-64-2P 609341-65-3P 609341-66-4P 609341-67-5P 609341-68-6P 609341-69-7P 609341-70-0P 609341-71-1P 609341-72-2P 609341-73-3P 609341-74-4P 609341-75-5P 609341-76-6P 609341-77-7P 609341-78-8P 609341-79-9P 609341-80-2P 609341-81-3P 609341-82-4P 609341-83-5P

RL: AGR (Agricultural use); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of (phenylethyl)acrylamides as agricultural fungicides) 609341-62-0 HCAPLUS

CN 2-Pyridineacetamide, N-[2-(3,4-dimethoxyphenyl)ethyl]- α -(2-methylpropylidene)-5-(trifluoromethyl)-, (α Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN

RN 609341-63-1 HCAPLUS

CN 2-Pyridineacetamide, N-[2-(4-ethoxy-3-methoxyphenyl)ethyl]- α -propylidene-, (α E)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 609341-64-2 HCAPLUS

CN 2-Pyridineacetamide, N-[2-(4-ethoxy-3-methoxyphenyl)ethyl]-5-methyl- α -propylidene-, (α Z)- (9CI) (CA INDEX NAME)

RN 609341-65-3 HCAPLUS

CN 2-Pyridineacetamide, N-[2-(4-ethoxy-3-methoxyphenyl)ethyl]- α -propylidene-, (α Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 609341-66-4 HCAPLUS

CN 2-Oxazoleacetamide, N-[2-(4-ethoxy-3-methoxyphenyl)ethyl]-4-methyl- α -propylidene-, (α Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 609341-67-5 HCAPLUS

CN 2-Pyridineacetamide, 5-bromo- α -(cyclohexylmethylene)-N-[2-(3,4-dimethoxyphenyl)ethyl]-, (α Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 609341-68-6 HCAPLUS

CN 2-Pyridineacetamide, N-[2-(4-ethoxy-3-methoxyphenyl)ethyl]- α -(2-methylpropylidene)-5-(trifluoromethyl)-, (α Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 609341-69-7 HCAPLUS

CN 2-Pyridineacetamide, 5-bromo- α -(cyclopropylmethylene)-N-[2-(3,4-dimethoxyphenyl)ethyl]-, (α Z)- (9CI) (CA INDEX NAME)

RN 609341-70-0 HCAPLUS

CN 2-Pyridineacetamide, α -(2,2-dimethylpropylidene)-N-[2-(4-ethoxy-3-methoxyphenyl)ethyl]-5-(trifluoromethyl)-, (α Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 609341-71-1 HCAPLUS

CN 2-Pyridineacetamide, N-[2-(3,4-dimethoxyphenyl)ethyl]- α -(2-ethylbutylidene)-5-(trifluoromethyl)-, (α Z)- (9CI) (CA INDEX NAME)

RN 609341-72-2 HCAPLUS

CN 2-Pyridineacetamide, N-[2-(3,4-dimethoxyphenyl)ethyl]- α -(2-ethylbutylidene)-5-methyl-, (α Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 609341-73-3 HCAPLUS

CN 2-Pyridineacetamide, N-[2-(3,4-dimethoxyphenyl)ethyl]- α -propylidene-5-(trifluoromethyl)-, (α Z)- (9CI) (CA INDEX NAME)

RN 609341-74-4 HCAPLUS

CN 2-Pyridineacetamide, 5-bromo-N-[2-(4-ethoxy-3-methoxyphenyl)ethyl]- α -propylidene-, (α Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 609341-75-5 HCAPLUS

CN 2-Pyridineacetamide, N-[2-(4-ethoxy-3-methoxyphenyl)ethyl]- α -propylidene-5-(trifluoromethyl)-, (α Z)- (9CI) (CA INDEX NAME)

RN 609341-76-6 HCAPLUS

CN 2-Pyridineacetamide, 5-bromo-N-[2-(3,4-dimethoxyphenyl)ethyl]- α -propylidene-, (α Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 609341-77-7 HCAPLUS

CN 2-Pyridineacetamide, 5-bromo-N-[2-(3,4-dimethoxyphenyl)ethyl]- α -(2-methylpropylidene)-, (α Z)- (9CI) (CA INDEX NAME)

RN 609341-78-8 HCAPLUS

CN 2-Pyridineacetamide, N-[2-(3,4-dimethoxyphenyl)ethyl]- α -(2,2-dimethylpropylidene)-5-(trifluoromethyl)-, (α Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 609341-79-9 HCAPLUS

CN 2-Pyridineacetamide, 5-bromo-N-[2-(3,4-dimethoxyphenyl)ethyl]- α -(2,2-dimethylpropylidene)-, (α Z)- (9CI) (CA INDEX NAME)

RN 609341-80-2 HCAPLUS

CN 2-Pyridineacetamide, α -(cyclopropylmethylene)-N-[2-(3,4-dimethoxyphenyl)ethyl]-5-(trifluoromethyl)-, (α Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 609341-81-3 HCAPLUS

CN 2-Pyridineacetamide, 5-bromo-N-[2-(3,4-dimethoxyphenyl)ethyl]- α -(2-methoxy-2-methylpropylidene)-, (α Z)- (9CI) (CA INDEX NAME)

RN 609341-82-4 HCAPLUS

CN 2-Pyridineacetamide, 5-bromo-N-[2-(3,4-dimethoxyphenyl)ethyl]- α -(2-methylbutylidene)-, (α Z)- (9CI) (CA INDEX NAME)

Double bond geometry as shown.

RN 609341-83-5 HCAPLUS

CN 2-Pyridineacetamide, 5-bromo-N-[2-(3,4-dimethoxyphenyl)ethyl]- α -(2-ethylbutylidene)-, (α Z)- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51 ANSWER 2 OF 8 HCAPLUS COPYRIGHT 2006 ACS on STN DUPLICATE 2

ACCESSION NUMBER:

2002:10427 HCAPLUS Full-text

DOCUMENT NUMBER:

136:69651

TITLE:

Preparation of acrylamide derivatives as agrochemical

fungicides

INVENTOR(S):

Sakaguchi, Hiroshi

PATENT ASSIGNEE(S):

Sumitomo Chemical Co., Ltd., Japan

SOURCE:

PCT Int. Appl., 96 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

Japanes

PATENT INFORMATION:

	PATENT NO.			KIND DATE			APPLICATION NO.					DATE						
	WO 2002000607		A1 20020103			WO 2001-JP5037					20010613							
		W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB	, BG,	BR,	BY,	BZ,	CA,	CH,	CN,
			CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC	, EE,	ES,	FI,	GB,	GD,	GE,	GH,
			GM,	HR,	HU,	ID,	IL,	IN,	IS,	ΚE,	KG	, KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,
			LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	ΜX	, MZ,	NO,	NZ,	PL,	PT,	RO,	RU,
			SD,	SE,	SG,	SI,	SK,	SL,	ТJ,	TM,	TR	, TT,	TZ,	UA,	UG,	US,	UZ,	VN,
			YU,	ZA,	ZW,	AM,	ΑZ,	BY,	KG,	KZ,	MD	, RU,	ТJ,	TM				
		RW:	GH,	GM,	KE,	LS,	MW,	MZ,	SD,	SL,	SZ	, TZ,	UG,	ZW,	AT,	BE,	CH,	CY,
			DE,	DK,	ES,	FI,	FR,	GB,	GR,	ΙE,	ΙT	, LU,	MC,	NL,	PT,	SE,	TR,	BF,
			ВJ,	CF,	CG,	CI,	CM,	GΑ,	GN,	GW,	ML	, MR,	NE,	SN,	TD,	TG		
	AU	2001	0642'	73		A 5	A5 20020108			AU 2001-64273					2	0010	613	
	ΕP	1295	868	•		A 1	A1 200303		0326	EP 2001-938646				20010613			613	
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR	, IT,	LI,	LU,	NL,	SE,	MC,	PT,
			IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL	, TR	•				-	
	JP	2002	3564	65		A2		2002	1213	JP 2001-187931				20010621			621	
	US	2003	1953	54		A1		2003	1016		US	2002-	3110	13		_ 2	0021	212
	US	6762	321			B2		2004	0713									
PRIOR	IT:	APP	LN.	INFO	.:						JΡ	2000-	1956	49		A 2	0000	629
							•				JP	2000-	3786	66		A 2	0001	213
											JΡ	2001-	9609	6		A 2	0010	329
										WO	2001-	JP50	37	1	w 2	0010	613	
ОПИБР	THE COURCE (C).				MAD	חתם	126.	COCE	1									

OTHER SOURCE(S):

MARPAT 136:69651

The title compds. I [R1 is C1-10 haloalkyl or the like; R2 is hydrogen or the AΒ like; X is oxygen or sulfur; Y is oxygen or sulfur; Ar is an aromatic group; A is ethylene or the like; and Z1 and Z2 are each alkyl, alkoxy, or the like] are prepared The title compound I [R1X = CH2FO; Ar = 4-methylphenyl; Y = O; R2 = H; A = CH2CH2; Z1 = Z2 = MeO] at 200 ppm gave 90% control of Plasmopara viticola.

384822-95-1P IT

> RL: AGR (Agricultural use); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of acrylamide derivs. as agrochem. fungicides)

384822-95-1 HCAPLUS RN

2-Thiopheneacetamide, α -[(difluoromethoxy)methylene]-N-[2-[3-methoxy-4-(2-propynyloxy)phenyl]ethyl]-4-methyl- (9CI) (CA INDEX NAME)

REFERENCE COUNT: THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L51. ANSWER 3 OF 8 MARPAT COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

143:248152 MARPAT Full-text

TITLE:

Preparation of amide derivative of Anona squamosa as

INVENTOR(S):

antiparkinsonian agents Liang, Xiaotian; Liu, Gengtao; Feng, Weihong; Ji,

Xiaoshen; Zhu, Liya; Xie, Ping; Wei, Huailing; Wang,

Qingli; Jiao, Xiaozhen

PATENT ASSIGNEE(S):

Institute of Materia Medica, Chinese Academy of

Medical Sciences, Peop. Rep. China

SOURCE:

Faming Zhuanli Shenqing Gongkai Shuomingshu, 46 pp.

CODEN: CNXXEV

DOCUMENT TYPE:

Patent

LANGUAGE:

Chinese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1445211	Α	20031001	CN 2002-107737	20020320
PRIORITY APPLN. INFO	. :		CN 2002-107737	20020320
OTHER SOURCE(S):	CA	SREACT 143:248	3152	
GI				

Title compds. represented by the formula I [wherein: ring A, B = (un)substituted Ph or aromatic heterocycle; R5 = H or alkyl or connected with the substituent of ring B by a covalent bond; R6 = H, alkyl, CO2H or ester group; n = 1-4; and their isomers thereof] were prepared as antiparkinsonian agents. For example, II was given in a multi-step synthesis starting from the reaction of 2,5-dimethoxybenzeneacetic acid with 4-acetoxy-3- methoxybenzene. II showed stimulation of movement recovery and increasing of learning ability in MPTP model rats action test, etc. Thus, I and their pharmaceutical compns. are useful for the prevention and treatment of Parkinson's diseases and Alzheimer's diseases, and improvement of the memory.

MSTR 1

G1 = heteroaryl <containing 1 or more heteroatoms>

(opt. substd. by 1 or more G3)

G2 = Ph (opt. substd. by 1 or more G3)

G3 = alkoxy <containing 1-7 C>

G5 = NH

G8 = (1-4) CH2

 $G9 = 1-14 \ 3-4$



Patent location:

claim 1

Note:

additional ring formation also claimed

ACCESSION NUMBER:

L51 ANSWER 4 OF 8 MARPAT COPYRIGHT 2006 ACS on STN

128:48241 MARPAT Full-text

TITLE:

Preparation of 3-(piperazinophenyl)acrylamides and

analogs as 5-HT1 receptor ligands

INVENTOR(S):

Howard, Harry Ralph; Segelstein, Barbara Eileen

PATENT ASSIGNEE(S):

Pfizer Inc., USA

SOURCE:

Eur. Pat. Appl., 29 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PAI	ENT NO	٠.	KIND	DATE		APPLI	CATION N	0.	DATE	*		
	EP	810220		A1	19971203		EP 19	97-30299	5	19970501			
	ΕP	810220		В1	20011212								
		R: A	T, BE	, CH, DE	, DK, ES,	FR,	GB, GR,	IT, LI,	LU,	NL, SE,	PT,	IE,	FI
	AT	210649		E	20011215		AT 19	97-30299	5	19970501			
	ES	216604	6	Т3	20020401		ES 19	97-30299	5	19970501			
	JP	100957	65	A2	19980414		JP 19	97-13080	0	19970521			
	JP	302694	8	B2	20000327								
	CA	220612	2	АĀ	19971128		CA 19	97-22061	22	19970526			
	CA	220612	2	С	20020305					,			
	US	625895	3	B1	20010710		US 19	97-86459	3	19970528			
PRIO	RITY	APPLN	. INF	0.:			US 19	96-18580	P	19960528			
											-	_	

ΑB R1ZCR2R6CR5R6CONR3R4 [I; R1 = e.g., 4-(un)substituted-1-piperazinyl wherein substituents may be alkyl, alkyl(hetero)aryl, etc.; R2 = H, alkyl, (un) substituted Ph, etc.; R3 = H, alkyl, phenyl(alkyl), etc.; R4 = alkyl or aryl; NR3R4 = heterocyclyl; R5 = H, alkyl, aryl; R6,R7 = H; R6R7 = bond; Z = (un) substituted 1,2-phenylene] were prepared Thus, 2-(4-methyl-1piperazinyl)benzaldehyde was condensed with PhCH2CONHPh to give 2-R1C6H4CH:CPhCONHPh (R1 = 4-Methyl-1-piperazinyl). Data for biol. activity of I were given.

MSTR 1

G25 = 229

296-G27

G26 = (1-3) CH2

G27 = Ph (opt. substd. by 1 or more G28)

G28 = alkoxy <containing 1-6 C>

G30 = 12

G29 12——G25

G32 = pyridyl (opt. substd.)

Derivative: or pharmaceutically acceptable salts

Patent location: claim 1

L51 ANSWER 5 OF 8 MARPAT COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 132:60481 MARPAT Full-text

TITLE: Fluorodipheny acrylamide-containing fungicide INVENTOR(S): Li, Zongcheng; Liu, Changling; Liu, Wucheng

PATENT ASSIGNEE(S): Shenyang Chem. Inst., Ministry of Chem. Industry,

Peop. Rep. China

Patent

SOURCE: Faming Zhuanli Shenqing Gongkai Shuomingshu, 25 pp.

CODEN: CNXXEV

DOCUMENT TYPE:

LANGUAGE: Chinese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
CN 1167568	A	19971217	CN 1996-115551	19960821
CN 1043720	В	19990623		
PRIORITY APPLN. INF	o.:	•	CN 1996-115551	19960821
			CN 1995-108849	19950828
GI				

$$R^{20}$$
 $C = C = C = N = ZR^4$

The acrylamide (I) (R1, and/or R2 = C1-6 alkyl, C1-6 haloalkyl, C3-6 AB cycloalkyl, C3-6 cycloalkyl-C1-6 alkyl etc.; R3 = H, -CN, imidazolyl, C3-6 alkyl etc.; X = 0, S, or NH; Z = bond or O; R4, and/or R5 = H, C1-6 alkyl, C2-6 alkenyl, C3-6 alkynyl etc.) has fungicidal activity and may be mixed with other known fungicide. The dosage form of the acrylamide is selected from emulsifiable solution, powder, wetting powder, suspension, and granule. The carrier for powder, wetting powder, and granule is selected from kieselguhr, clay, gypsum, talc, and kaolin; the solvent for emulsifiable solution from benzene, toluene, xylene, alkylbenzene, C1-6 fatty alc., benzenemethanol, cyclohexanol, acetone, butanone, Me iso-Bu ketone, DMF, DMSO, Nmethylpyrrolidone, water, etc. The fungicide may contain surfactant as emulsifier, dispersant, or wetting agent. The surfactant is selected from sodium laurylbenzenesulfonate, K- 12, polyoxyethylene fatty acid ester, polyoxyethylene fatty acid alc., polyoxyethylene fatty acid amine, ethoxycastor oil, sodium lignosulfonate, carboxymethyl alc., polyvinyl alc., and polyvinyl ester.

MSTR 1

G4 = triazolyl G5 = O G6 = 71

79-G10

G7 = alkoxy G8 = Ph (opt. substd. by (1-5) G7) G9 = NH

G10 = 79

7613—G8

G13 = alkylene <containing 1-3 C> Patent location: claim 1

L51 ANSWER 6 OF 8 MARPAT COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 132:9930 MARPAT Full-text

TITLE:

Acrylamide germicide containing fluoro-diphenyl group

INVENTOR(S):

Li, Zongcheng; Liu, Changling; Liu, Wucheng Shenyang Chemical Inst., Ministry of Chemical

PATENT ASSIGNEE(S):

Industry, Peop. Rep. China

SOURCE:

Faming Zhuanli Shenqing Gongkai Shuomingshu, 25 pp.

CODEN: CNXXEV

DOCUMENT TYPE:

Patent

LANGUAGE:

Chinese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

kЗ

I

The compound I [R1, or/and R2 = C1-6 alkyl, C3-6 cycloalkyl, C2-6 alkenyl, C2-6 alkynyl, aryl etc.; R3 = H, CN, NO2, triazolyl, pyridyl, imidazolyl, C1-6 alkyl etc.; X = O, S, or NH; Z = O, or O; R4, or/and R5 = H, C1-6 alkyl, C2-6 alkenyl, C2-6 alkynyl, C1-6 alkylcarbonyl, C3-6 cycloalkyl etc.] is germicidal and may be used by mixing with other known germicides. The dosage form may be emulsifiable solution, powder, wetting powder, suspensoid, and granule. The carrier is selected from zeolite, clay, gypsum, talc, and kaolin; the solvent from benzene, toluene, xylene, alkylbenzene, benzyl alc., cyclohexanol, acetone, butanone, Me iso-Bu ketone, DMF, DMSO, N-methylpyrrolidone, and water etc.; and the surfactant from K-12, Na lauryl benzene sulfonate, polyvinyl fatty acid ester, polyvinyl fatty acid alc., polyvinyl fatty acid amine, ethoxy castor oil, Na or K lignosulfonate, carboxylmethyl alc., polyvinyl alc., and polyvinyl ester.

MSTR 1

G4 = triazolyl

G5 = O G6 = 71

7G9—G10

G7 = alkoxy

G8 = Ph (opt. substd. by (1-5) G7)

G9 = NH G10 = 79

7G13—G8

G13 = alkylene <containing 1-3 C>
Patent location: claim 1

L51 ANSWER 7 OF 8 MARPAT COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

125:114317 MARPAT <u>Full-text</u> Fungicidal carboxamides

TITLE: INVENTOR(S):

Seitz, Thomas; Heinemann, Ulrich; Stenzel, Klaus;

Dutzmann, Stefan

PATENT ASSIGNEE(S):

Bayer A.-G., Germany

SOURCE:

Ger. Offen., 29 pp.

CODEN: GWXXBX

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

1

PATENT INFORMATION:

_	PAT	CENT	NO.		KII	ND.	DATE			A)	PPLI	CATI	ои ис	ο.	DATE			•
												<u></u> -						
	DE 4443641 A1			1	19960613			DE 1994-4443641				41	19941208			•		
	WO 9617825 A1			1	1996	0613		WO 1995-EP4668				В	19951127					
		W:	AU,	BB,	BG,	BR,	BY,	CA,	CN,	CZ,	FI,	HU,	JP,	KR,	ΚZ,	LK,	MX,	NO,
			NZ,	PL,	RO,	RU,	SK,	UA,	US									
		RW:	ΑT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE,
			BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	ML,	MR,	NE,	SN,	TD,	TG		
	AU	9642	570		A:	1	1996	0626		Α	J 19	96-4	2570		1995	1127		
	EP	7962	42		A.	1	1997	0924		F.3	P 19	95-9	4102	9	1995	1127		

R: BE, CH, DE, FR, GB, IT, LI, NL

19990106 JP 11500103 T2

JP 1995-517296 19951127 19941208 DE 1994-4443641

PRIORITY APPLN. INFO .: 19951127 WO 1995-EP4668

GΙ

Carboxamides, such as I [X = bond, CH2, CHMe, CMe2; R = substituted Ph] were AΒ prepared Thus, the acid was amidated with PhCH2NH2 to give I [X = CH2, R = Ph] which at 250 g/ha protected barley against Erysiphe graminis.

MSTR 1

$$G2 = 10$$

$$G35 = NH$$

$$G40 = 372$$

Patent location:

claim 1

L51 ANSWER 8 OF 8 MARPAT COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

114:37800 MARPAT Full-text

TITLE: INVENTOR(S): Preparation of N-benzylcarboxamides as herbicides. Oba, Nobuyuki; Sato, Masahiro; Ikeda, Atsuhiko; Takeuchi, Akira; Matsunari, Kenji; Yamada, Yuji;

Nakamura, Michiya; Nakamura, Yasuo

PATENT ASSIGNEE(S):

Kumiai Chemical Industry Co., Ltd., Japan; Ihara

Chemical Industry Co., Ltd.

SOURCE:

Jpn. Kokai Tokkyo Koho, 11 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE		
JP 02200658	A2	19900808	JP 1989-18777	19890128		
PRIORITY APPLN. INFO.	:		JP 1989-18777	19890128		
GI						

Herbicides contain the title compds. I [R1 = 1-naphthyl, (halo- or Mesubstituted) Ph or thienyl; R2 = H, Me; R3 = H, Cl, lower alkyl, OH, MeO; R4, R5 = Me, Et; R4R5, = cyclopropylidene; X = halo, Me, MeO, PhO, CF3, CO2Et; Y = N, CH; m = 0-3; n = 0, 1] as active ingredients. (E)-2-Phenyl-2-butenoyl chloride in acetone was treated with α -ethyl- α -methylbenzylamine and NaHCO3 at room temperature for 3 h to give 81% (E)-I (R1 = Ph, R3 = R4 = Me, R5 = Et, Xm = H, Y = CH, n = 0), which (100 g/10 are) showed $\geq 90\%$ herbicidal effect against Echinochloa crus-galli, Cyperus difformis, Monochoria vaginalis, and Scirpus juncoides with $\leq 10\%$ damage on rice.

MSTR 1

$$G1 = bond$$

$$G2 = 11$$

$$G3 = 22 / 23 / 26$$

$$G8 = 45$$

$$G9 = Me$$
 $G10 = OMe$

Patent location:

claim 1

INVENTOR SEARCH

=> fil hcap medline embase biosis

FILE 'HCAPLUS' ENTERED AT 12:00:22 ON 05 DEC 2006

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=> d que 134 L17228 SEA GRAMMENOS W?/AU L18 380 SEA GROTE T?/AU L19 77 SEA BLETTNER C?/AU L20 148 SEA GEWEHR M?/AU 120 SEA GYPSER A?/AU L21 5036 SEA MULLER B?/AU L22 L23 253 SEA RHEINHEIMER J?/AU 691 SEA SCHAFER P?/AU L24 L25 11 SEA SCHWOGLER A?/AU L26 275 SEA TORMO J?/AU 99 SEA GOTZ N?/AU L27 L28 1238 SEA LORENZ G?/AU L29 842 SEA AMMERMANN E?/AU L30 479 SEA STRATHMANN S?/AU L31 235 SEA STIERL R?/AU L34 2 SEA (L17 OR L18 OR L19 OR L20 OR L21 OR L22 OR L23 OR L24 OR L25 OR L26 OR L27 OR L28 OR L29 OR L30 OR L31) AND PHENETHY? AND ?ACRYLAMID?

=> d 134 ibib abs hit 1-2

L34 ANSWER 1 OF 2 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2001:923549 HCAPLUS Full-text

DOCUMENT NUMBER:

136:33328 .

TITLE:

Preparation of phenethylacrylamides as

INVENTOR(S):

fungicides

Grammenos, Wassilios; Sauter, Hubert;
Cullmann, Oliver; Gewehr, Markus; Mueller,

Bernd; Tormo i Blasco, Jordi; Goetz, Norbert; Volk,

Thorsten; Lorenz, Gisela; Ammermann,

Eberhard; Stierl, Reinhard;

Strathmann, Siegfried

PATENT ASSIGNEE(S):

Basf Aktiengesellschaft, Germany

SOURCE:

PCT Int. Appl., 44 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent German

LANGUAGE: GEFAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.

KIND DATE

APPLICATION NO.

DATE

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WO 2001095721
                                20011220
                                             WO 2001-EP6686
                                                                    20010613
                          A2
             AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH,
             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT,
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             UZ, VN, YU, ZA, ZW
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
     CA 2412489
                          AΑ
                                20021211
                                            CA 2001-2412489
                                                                    20010613
                                             EP 2001-964978
     EP 1289365
                          A2
                                20030312
                                                                    20010613
     EP 1289365
                          В1
                                20040908
            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
     BR 2001011611
                                20030701
                                             BR 2001-11611
                          Α
                                                                    20010613
     HU 200300650
                          A2
                                20030728
                                             HU 2003-650
                                                                    20010613
     JP. 2004503475
                          T2
                                20040205
                                             JP 2002-509917
                                                                    20010613
     NZ 522952
                          Α
                                20040430
                                             NZ 2001-522952
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     EE 200200685
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                                20040615
                                             EE 2002-685
                                                                    20010613
     AT 275340
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                                20040915
                                             AT 2001-964978
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     ES 2227256
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                                20050401
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                          A1
                                             US 2002-297287
     US 2003191190
                                20031009
                                                                    20021204
    US 6696607
                          В2
                                20040224
     BG 107358
                          Α
                                20030731
                                             BG 2002-107358
                                                                    20021205
     ZA 2003000322
                          Α
                                20040121
                                             ZA 2003-322
                                                                    20030113
PRIORITY APPLN. INFO.:
                                             DE 2000-10028576
                                                                 Α
                                                                    20000614
                                             DE 2000-10028857
                                                                 A 20000614
                                             WO 2001-EP6686
                                                                 W 20010613
OTHER SOURCE(S):
                         MARPAT 136:33328
```

GΙ

AB The **phenethylacrylamides** I [X = halo, alkyl, haloalkyl, alkoxyhaloalkoxy, etc.; m, n = 1-4; Y = halo, nitro, cyano, alkyl, CF3, alkoxy or phenyl; R1, R2 = H, halo, alkyl, alkoxy, haloalkoxy or CF3; R3, R4, R5, R6 = H, alkyl or alkoxy; R3CR4 = cyclopropyl] are prepared as fungicides.

TI Preparation of phenethylacrylamides as fungicides

IN Grammenos, Wassilios; Sauter, Hubert; Cullmann, Oliver;
Gewehr, Markus; Mueller, Bernd; Tormo i Blasco, Jordi; Goetz,
Norbert; Volk, Thorsten; Lorenz, Gisela; Ammermann,
Eberhard; Stierl, Reinhard; Strathmann, Siegfried

AB The **phenethylacrylamides** I [X = halo, alkyl, haloalkyl, alkoxyhaloalkoxy, etc.; m, n = 1-4; Y = halo, nitro, cyano, alkyl, CF3, alkoxy or phenyl; R1, R2 = H, halo, alkyl, alkoxy, haloalkoxy or CF3; R3, R4, R5, R6 = H, alkyl or alkoxy; R3CR4 = cyclopropyl] are prepared as fungicides.

ST **phenethylacrylamide** deriv prepn fungicide

```
Fungicides
ΙT
        (agrochem.; phenethylacrylamide derivs.)
IT
     554-52-9P, 4-(2-Aminoethyl)-2-methoxyphenol
                                                   24091-92-7P
                                                                  37542-28-2P
     82549-10-8P, 3,3-Dichloro-2-(4-chlorophenyl)acrylic acid
                                                                380610-20-8P,
     3,3-Dichloro-2-(4-chlorophenyl)-acrylic acid methyl ester
                                                                  380610-21-9P
     RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (intermediate in preparation of phenethylacrylamide fungicide)
ΙT
     106-96-7, Propargyl bromide
                                   120-20-7, Homoveratrylamine
     52449-43-1, Methyl 4-chlorophenylacetate
     RL: RCT (Reactant); RACT (Reactant or reagent)
        (reactant in preparation of phenethylacrylamide fungicide)
L34 ANSWER 2 OF 2
                    BIOSIS COPYRIGHT (c) 2006 The Thomson Corporation on STN
ACCESSION NUMBER:
                    2004:168854 BIOSIS
                                         Full-text
DOCUMENT NUMBER:
                    PREV200400170720
TITLE:
                    Use of phenethyl acrylamides, novel
                    phenethyl acrylamides, method for the
                    production thereof and agents containing the same.
                    Grammenos, Wassilios [Inventor, Reprint Author];
AUTHOR(S):
                    Sauter, Hubert [Inventor]; Cullmann, Oliver [Inventor];
                    Gewehr, Markus [Inventor]; Muller, Bernd
                    [Inventor]; Blasco, Jordi Tormo i [Inventor]; Gotz,
                    Norbert [Inventor]; Volk, Thorsten [Inventor];
                    Lorenz, Gisela [Inventor]; Ammermann,
                    Eberhard [Inventor]; Stierl, Reinhard
                    [Inventor]; Strathmann, Siegfried [Inventor]
CORPORATE SOURCE:
                    Ludwigshafen, Germany
                    ASSIGNEE: BASF Aktiengesellschaft, Ludwigshafen, Germany
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AB
     Use of phenethylacrylamides of the formula I: ##STR1## in which the
     substituents have the following meanings: X is halogen, alkyl, haloalkyl,
     alkoxy, haloalkoxy and --O--C(Ra,Rb)--CidentC--R6; Ra,Rb and Rc have the
     meanings given in the description; m,n independently of one another are 1 to
     4, it being possible for the radicals X or Y to be different if m or n is
     greater than 1; Y is halogen, nitro, cyano, alkyl, CF3, alkoxy and phenyl;
     R1,R2 independently of one another are hydrogen, halogen, alkyl, alkoxy,
     haloalkoxy and CF3; R3,R4,R5,R6 independently of one another are hydrogen,
     halogen, alkyl, alkoxy, or R3 and R4 together form a cyclopropyl ring, it
     being possible for the C--R5 -- and C--R6 bonds can be in the E- or Z-position
     relative to each other; for controlling phytopathogenic fungal pests, novel
     phenethylacrylamides, their preparation, and compositions comprising them.
TI
     Use of phenethyl acrylamides, novel phenethyl
     acrylamides, method for the production thereof and agents
     containing the same.
ΑU
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     Siegfried [Inventor]
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Use of phenethylacrylamides of the formula I: ##STR1## in which the substituents have the following meanings: X is halogen, alkyl, haloalkyl, alkoxy, haloalkoxy and --O--C(Ra,Rb)--CidentC--R6; Ra,Rb and Rc have the meanings given in the description; m,n independently of one another are 1 to 4, it being possible for the radicals X or Y to be different if m or n is greater than 1; Y is halogen, nitro, cyano, alkyl, CF3, alkoxy and phenyl; R1,R2 independently of one another are hydrogen, halogen, alkyl, alkoxy, haloalkoxy and CF3; R3,R4,R5,R6 independently of one another are hydrogen, halogen, alkyl, alkoxy, or R3 and R4 together form a cyclopropyl ring, it being possible for the C--R5 -- and C--R6 bonds can be in the E- or Z-position relative to each other; for controlling phytopathogenic fungal pests, novel phenethylacrylamides, their preparation, and compositions comprising them.

IT Major Concepts

Pharmaceuticals (Pharmacology)

IT Chemicals & Biochemicals

phenethyl acrylamides: pharmaceutical